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Description

This invention relates to coated ophthalmic lenses and to methods for applying coatings to such lenses. More particularly it relates to cast opthalmic lenses, such as spectacle lenses, which have an optically clear, aberration free, abrasion-resistant coating applied thereto and to the methods of applying a coating by the application of a thin layer of a composition containing reactive ethylenically-unsaturated groups.

The application of coatings to surfaces is known. For example, U.S. Patents 4,073,967 and 4,084,021 disclose the application of abrasion-resistant coatings to surfaces by spin coating, dip coating, spray coating, and flow coating techniques.

U.S. 4,319,811 discloses abrasion-resistant radiation-curable coatings which are the polymerisation products of compositions containing reactive ethylenically unsaturated groups. These coatings are applied by techniques such as spraying, brushing, dipping and roll coating to plastic lenses such as the lenses of suppliesses, for example.

While these techniques have been successful when used to coat smooth surfaces, such as the optical surface of single vision spectacle lenses, they have not been entirely satisfactory. For example, such techniques result in a build-up of a coating material at the outer edge of the lens which causes optical aberrations. Such techniques are even less satisfactory when they are used to coat irregular surfaces, such as those present on multivision ophthalmic devices, particularly those made from plastic. Such devices generally have segments which project abruptly away from (e.g. project above or below) at least one major optical surface of the device. Coating solutions applied to these optical surfaces by spin, dip, spray, or flow techniques must flow around and over these segments. The resulting coating generally exhibits flow marks around the projection and is thick as the intersections of the segment and the optical surface. This intersection is referred to hereinafter as the segment line.

Vapor coating techniques may also be employed. However, the resulting coatings tand to be very thin, for example 0.5 micron thick, and consequently are not useful in providing significant abrasion resistance.

U.S. Patent 4,081,578 discloses a molding method wherein parts are first cast in a mold after which the mold is opened to provide a space for the injection of a predetermined quantity of coating composition. The mold is closed to spread the coating composition over the surface of the cast part, the coating cured in the closed mold, and the coated part removed from the mold. This technique is complicated and does not provide good adhesion of the coating to the substrate surface.

In yet another technique a gel coating material is applied to the surface of a mold and allowed to partially cure. See U.S. Patent 4,081,578 at col. 2, lines 15—22. A mixture of resin compound and glass fibers is applied to and rolled over the gel coat. The resulting composite structure is allowed to cure. The purpose of this technique is to fill in any defects on the surface of an article.

U.S. Patent 4,338,269 discloses a technique for applying a coating to the surface of a molded article. In this technique a composition containing a pertaerythritol base polyacrylate or polymethacrylate and a cellulose ester or a vinyl chloride-vinyl acetate containing polymer is applied to a mold and fully cured by exposure to actinic radiation. Subsequently a polymerizable composition is charged to the mold and polymerizad. Such compositions exhibit only minimal abrasion resistance and have not been found to provide optically clear coatings.

U.S. Patent 3,968,305 discloses a shaped article and a method for applying a "mar-resistant" layer thereto. One method disclosed in this patent involves the steps of first, applying to a mold a polymerizable material selected from

a) a compound having a total of at least three acryloxy and/or methacryloxy groups in the molecule which are linked with an aliphatic hydrocarbon residue having not more than 20 carbon atoms,

b) a monomer mixture comprising at least 20% by weight of said compound and at most 80% by weight of at least one other copolymerizable monomer, or

c) a partially polymerized product of said compound or said monomer mixture:

second, polymerizing the polymerizable material to form an abrasion-resistant film on the mold; third, charging a monomer to the mold; and

fourth, polymerizing the monomer,

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This method may be applied to the manufacture of coated sunglasses, fashion glasses and other glasses for example.

The ophthalmic lens and method of the present invention do not require the formation of an abrasionresistant film on the mold prior to introduction of the monomer. To the contrary such a technique would not work in the invention as is shown.

The present invention sims to provide an ophthalmic lens having a thin, optically clear, aberration free, abrasion-resistant coating on one or more of its optical surfaces. It further aims to provide a method of applying a coating to ophthalmic devices such as spectacle lenses, which is equally useful in applying a coating to and multifocal lenses.

The invention provides a method of providing a thin, optically clear coating to at least one optical surface of an ophthalmic lens comprising the steps of:

(i) applying a layer of a composition comprising a material containing reactive ethylenically unsaturated groups to at least a first face of a mold used to manufacture said ophthalmic lens, wherein said

face imparts a desired optical configuration to a first optical surface of said ophthalmic iens;

(ii) reacting said composition to a degree that said composition forms a film that is dry, non-abrasion resistant, at least weakly adhered to said first face, and exactly replicates said first face in an aberration-free manner such that said composition exhibits a degree of unsaturation in the range of 40% to 90% of the unsaturation it possessed prior to reaction;

(iii) filling said mod with an organic liquid material capable of hardening to a solid, room temperature-

stable state; and

(Iv) hardening said organic liquid material so as to form said ophthalmic lens, intimately bond said film to the hardened organic material, and adhere said film to said optical surface of said hardened organic material more firmly than it adheres to sald face, and

(v) post reacting said composition after said organic liquid has hardened to harden said composition. Preferably said material is a poly(ethylenically unsaturated alkoxyalkyl) compound of the formula -Z—A² wherein

(a) A¹ and A² are independently groups having terminal ethylenic unsaturation said groups having the

15 formula:

in which (1) R1 is selected from hydrogen or methyl

(2) R—O— is a monovalent residue of an aliphatic terminally unsaturated primary alcohol, ROH, where R is selected from the formulae:

so and

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wherein E is selected from the formulae:

a and c are independently integers of from 1 to 6, b is zero or an integer of from 1 to 6, R4 is selected from hydrogen or methyl, R⁵ is an eliphatic group having 1 to 15 carbon atoms which may be interrupted with up to two groups selected from the class consisting of ether or oxygen or

R⁵ having a valence of m+1 wherein m is an integer of 1 to 5,

(3) R2 is selected from the group of hydrogen,

wherein R⁶ is selected from alkyl and alkenyl groups having up to 5 carbon atoms, R⁷ is an aliphatic group of up to 8 carbon atoms or an eromatic group of up to 8 carbon atoms,

(4) R3 is an alkylene group having 1 to 8 carbon atoms and up to one caternary oxygen in the group, and (b) Z is a heterocyclic group of the formula:

wherein X is a divalent group required to complete a 5-membered heterocyclic ring and is selected from the group of:

wherein R⁸, R⁹, R¹⁰, and R¹¹ are independently selected from hydrogen, alkyl groups of 1 to 4 carbon atoms, cycloalky groups of 3 to 6 carbon atoms, and phenyl groups, and A³ is the same as A¹ and wherein said non-abrasion-resistant film is reacted in step (v) to harden it to an SEB abrasion resistance of at least 137.9 kPa (20 p.s.i.).

As used throughout this specification, the following terms have the following meanings:

(a) "Optically clear" means free from haze or other interference which prevents an object on one side of a lens from being seen clearly by an observer on the other side of a lens,

(b) "Aberration-free" means that an object on one side of a layer of the coated lens does not appear to be bent, twisted or distorted to a viewer on the other side of the lens.

(c) "Reactive ethylenically unsaturated groups" are those groups which polymerize, crosslink, or both when subjected to activating conditions described hereinafter.

(d) "Optical surface" means one of the surfaces which provides optical correction.

(e) "SEB abrasion resistance" is the abrasion resistance according to the technique described in US—A—4,084,091. Therefore the value reported for SEB abrasion resistance refers to the level of force necessary to cause visible scratching on the coated surface of the lens. Coatings applied by the process of the invention are thin (typically having a thickness of from about 0.5 to 50, and preferably of from about 2 to 10, and more preferably of from about 3 to 5, (microns µm thick). Furthermore, the coatings are free from surface aberrations due to nonuniformity of the thickness of the coating solution, flow marks, coating buildup, particularly at the segment lines and edges of the ophthalmic devices. Additionally, they exactly replicate the mold surface from which the ophthalmic device is made and consequently provide an optical surface having the desired surface configuration.

DETAILED DESCRIPTION

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Molds used in the manufacture of ophthalmic lenses from organic materials are generally made from glass or metal and typically have first and second mold sections which form the front and back optical surfaces on the devices. At least one of these sections has a surface which forms a finished optical surface.

The coating composition may be applied to one or more of the mold surfaces by a variety of techniques including spraying, dipping, brushing, flow coating, spin coating and the like. Flow coating has proven to be a very useful technique. In this technique a mold is preferably positioned such that the surface to be coated is oriented at an angle of from 45° to 60° from the horizontal. If the mold is designed for the production of multifocal devices having a straight segment line (such as a "flat top" lens), the face containing the multifocal portion is positioned so that the straight segment line is parallel to the vertical. In this position the outline of the multifocal portion of a common flat top bifocal lens mold forms the letter """

The molds are then preferably cleaned of particulate material by, for example, directing a stream of filtered nitrogen gas across the surface to be coated. A predetermined quantity of the composition is then applied to the top edge of the first face of the mold, preferably across an arc having a central angle of from about 140° to 160°. The composition is allowed to flow down to the bottom of the lens where any excess may be removed by an absorbent material placed in close proximity to the bottom of the die. If the composition includes a solvent, it may be, and preferably, is removed. This may be done by the use of infrared radiation or heat.

The coating is then reacted to a degree that it forms a dry film on the mold which exactly replicates the mold face in an aberration-free manner. This may be done by either heating the composition for a time sufficient to form the dry film or by exposing it to activating radiation for a time sufficient to form the dry film. Whichever technique is employed, it is essential that the coating be partially reacted within certain limits. For example, if the coating is underreacted, it provides a hazy film and may also exhibit poor abresion resistance. If, however the coating is overreacted, it exhibits unacceptable adhesion.

When thermal techniques are employed, the coating can be successfully partially reacted by exposing it to temperatures in the range of about 35° to 100°C for from about one to ten minutes.

When activating radiation, such as ultraviolet light or electron beam radiation, is used, the coating is reacted until it exhibits a degree of unsaturation of from about 40% to 90% of the unsaturation it possessed prior to reaction. Preferably it is reacted until it exhibits a degree of unsaturation in the range of from about 40% to 75%, and most preferably from about 55% to 70%, of the unsaturation it possessed prior to reaction. Coatings which have a degree of unsaturation above about 90% provide a hazy film (i.e., one which is not optically clear) on the final device and may also exhibit poor abrasion-resistance while those which have a degree of unsaturation below about 30% exhibit unacceptable adhesion to the final product.

After being partially reacted, the coating is dry, although it may be tacky to the touch. Additionally, the

coating will not flow by itself although it can be deformed by touching it with finger pressure. Furthermore, the coating is not abrasion-resistant at this point.

The specific technique used to bring about partial reaction may be selected to suit the nature of the coating composition. For example, when the reaction ethylenically unsaturated material contains acrylates functional groups, a satisfactory level of reaction can be achieved by exposing the coating composition to ultraviolet light. While the exact exposure conditions employed are dependent upon the intensity of the light, the distance of the coating from the light, and final degree of unsaturation desired, satisfactory partial reaction may be achieved by positioning the coating at a distance of from 1 to 90 cm from the light and exposing it to rediction from a 40 watt/cm Hanovia lamp for from 0.02 sec to 40 mln. Generally, the closer the coating is to the light source, the shorter the time of exposure.

If the ethylenically unsaturated materials contain alkoxy silene, the coatings can be satisfactorily reacted by the thermal techniques described above. Other techniques suitable for reacting the coatings will

be obvious to those skilled in the art.

After the desired level of reaction is obtained, the mold is assembled and filled with liquid organic material to provide the substrate. A wide veriety of thermosetting and thermoplastic organic materials can be employed. All that is necessary of such materials is that they be capable of solidifying to form a room-temperature stable, optically clear ophthalmic device. Examples of useful thermosetting materials include allyl diglycol carbonate monomer, acrylate monomers, and acrylate oligomers. Examples of useful thermoplastic materials include polycarbonate, polymethylmethacrylate, polyvinyl chloride, and the like.

Once the mold is filled, the organic liquid is hardened by any suitable technique. For example, allyl diglycol carbonate may be hardened by subjecting it to heat in the range of 35°C to 65°C for up to 16 hours

in the presence of a suitable polymerization agent.

As the organic liquid hardens, the film of coating composition forms an intimate bond to the substrate so that it adheres to the optical surface of the device more firmly than it adheres to the face of the mold. The coating is so intimately attached to the substrate that it can withstand repeated exposure to freeze/thew cycles without any loss of adhesion. This is characteristic of a chemical bond between the two materials.

Additionally as the organic liquid hardens the coating is further reacted so that once hardening of the

substrate is completed the coating has been rendered abrasion-resistant.

Once hardened the cast ophthalmic devices are removed from the mold. As the adhesion of the coating is greater to the surface of the ophthalmic device than it is to the face of the mold, it separates essentially completely from the mold. The resulting coating is thin and its exposed surface (that is the surface that formerly contacted the mold) provides exactly the optical surface desired on the ophthalmic device. Furthermore, it is free from aberrations such as would be caused by flow lines and buildup at segment lines which may be present.

The coated device may be used as is or alternatively it may be post reacted. Post-reaction may be accomplished by the same techniques utilized to initially react the coating. Typically the conditions employed during post curing are sufficient to assure essentially complete reaction of the coating. Surprisingly, post curing, even to a level such that there is essentially no available residual unsaturation

remaining, does not adversely affect the adhesion of the coating to the cast article.

While the previous discussion has been directed to the application of the coating to only one surface of a multifecal ophthalmic device, the process may also be used to apply the coating to single vision lenses, particularly those having a high curvature, and to two or more surfaces of an ophthalmic device.

Coetings produced by the method of the invention typically exhibit an SEB abrasion resistance of at least 137.9 kPa (20 psi). This means that a force of at least 137.9 kPa (20 psi) must be applied to the coating to cause visible scratching. This level of abrasion resistance represents a substantial improvement over the abrasion resistance of uncoated lenses. For example lenses made from allyl diglycol carbonate monomer scratch visibly after the application of only 6.89 to 20.67 kPa (1 to 3 psi). Although an abrasion resistance of (20 psi) 135 kPa represents a substantial improvement over this, it is only a fair or acceptable level of abrasion resistance in the context of the present invention. Thus, the lenses of the invention preferably exhibit an abrasion resistance of at least 275 kPa (40 psi) (a good level of abrasion resistance) and more preferably exhibit an abrasion resistance of at least 405 kPa (60 psi) (an excellent level of abrasion resistance).

The coating compositions useful in the method of the invention contain at least one material having reactive ethylenically unsaturated groups. Example of such materials are acylates, methacrylates, acrylic anhydrides, ethylenically unsaturated anhydrides, olefinic compounds, acrylamides, ethylenically unsaturated amides and urethanes, vinyl ester, vinyl ethers, vinyl halides, vinyl epoxy resins, vinyl silanes and siloxanes, vinyl heterocycles, melamines, and prepolymers and polymers of these materials. These materials may be mono- or polyfunctional.

Specific examples of these materials include methyl methacrylate, ethyl acrylate, 2-ethylhexyl acrylate, cyclohexyl acrylate, styrene, butadlene, 2-chlorostyrene, 2,4-dichlorostyrene, acrylic acid, acrylamide, acrylonitrile, t-butyl acrylate, methyl acrylate, butyl acrylate, N-vinyl pyrrolidone, 2-(N-butylcarbamyl)ethyl methacrylate, and 2-(N-ethylcarbamyl)ethyl methacrylate. Examples of other useful meterials include 1,4-butylene dimethacrylate or acrylate, ethylene dimethacrylate, hexanedioi diacrylate or dimethacrylate, glyceryl diacrylate or trimethacrylate, tripropylene glycol diacrylate, glyceryl triacrylate or trimethacrylate, pentaerythritol tetreacrylate or

tetramethacrylate, dimethiolpropane tetracrylate, dielilyi phthalate, dipentaerythritol pentaacrylate, dipentaerythritol hydroxypentaacrylate, neopantylgiycol triacrylate and 1,3,5-tri(2-methacryloxysthyl)-striazine, hexamethoxymethyl melamine, poly(acrylic acid) a poly(methacrylic acid), poly(methyl methacrylate), poly(vinyl chloride), poly(vinyl acatate, poly(vinyl butyral) and the like.

A specific class of useful materials comprises the polyacrylate functional hydrocarbons having at least two acrylate groups per molecule. A subclass of these materials which are specified for the coating compositions of ophthalmic lenses in accordance with the invention are the polylethylenically unsaturated alkoxyalkyl) heterocyclic compounds having the general formula

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(a) A¹ and A² Independently are alkoxyalkyl gorups having terminal ethylenic unsaturation and having the general formula

in which

(1) R1 is selected from hydrogen or methyl

(2) R—O— is a monovalent residue (formed by removal of the active hydrogen from an —OH group) of an aliphatic terminally unsaturated primary alcohol. ROH, R having the formula

or preferably

wherein E is selected from

a and c are independently an integer of from 1 to 6, b is zero or an integer of 1 to 6, R⁴ is selected from hydrogen or methyl, R⁵ is an aliphatic group having 1 to 15 carbon atoms (preferably alkylene of up to 15 carbon atoms) and optionally one or two caternary (i.e., backbone) oxygen atoms or

R5 having a valence of m+1, and m is an integer of 1 to 5,

(3) R2 is preferably hydrogen but can be

wherein R⁶ is preferably alkenyl but can be alkyl (each preferably having 2 to 5 carbon atoms) and can be substituted by a phenyl or carboxyl group, and R⁷ is an aliphatic group (of up to eight carbon atoms, e.g., alkyl) or aromatic group (preferably having up to 8 carbon atoms and more preferably a phenyl group) and R⁷ is most preferably an acryloyloxyalkyl or a methacryloyloxyalkyl group;

(4) R3 is an alkylene group having 1 to 6 carbon atoms and optionally one caternary oxygen atom; and

(b) Z is a heterocyclic group of the formula:

65 wherein X is a divalent group which is required to complete a 5- or 6-membered heterocyclic ring.

Preferably X is

but X can be

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wherein R^a, R^a, R¹⁰, and R¹¹ are independently hydrogen or lower alkyl (of 1 to 4 carbon atoms), cycloalkyl 15 (of 3 to 6 carbon atoms) or phenyl group (of 6 to 12 carbon atoms), and A³ is an alkoxyalkyl group as defined above for A¹ and A².

A specific heterocyclic compound of the generic formula is

wherein R12 Is

Materials of the formula A¹—Z—A² are disclosed in U.S. Patent 4,306,954.

Other polyacrylate functional materials useful in the method of the invention include "Sartomer" 295 (pentaerythritol tetraacrylate) and "Sartomer" 399 (dipentaerythritol monohydroxy pentaacrylate) both commercially available from Sartomer Company. These materials respectively have the formulae

CH2=CHCOCH2 CH2=CHCOCH2CCH2CH=CH2

Other specific classes of useful ethylenically unsaturated materials useful in the method of the invention include vinyl silene and siloxene materials and acryloxysllenes and siloxenes. Examples of such materials may be represented by the formula:

55 wherein R13 is selected from a vinyl group, e.g.,

or a methacryloxy group, e.g.,

CH_a=CH

CH_a=C C-O-R¹⁸

wherein R¹⁸ is an alkylene group of 1 to 8 carbon atoms, and R¹⁹ is hydrogen or hydrocarbyl radical of 1 to 8 carbon atoms (preferably alkyl). An epoxy group, such as one having of the following formulae

$$CH_2$$
 $CH-R_0^{17}$ or $CH-R_0^{17}$

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may be included in the silane or siloxane as the R19 group provided that it does not constitute all of the R13 groups. In these groups, R17 is a divalent hydrocarbon radical (allphatic-, aromatic-, or aliphatic- and aromatic-containing) of less than 20 carbon atoms or a divalent radical of less than 20 carbon atoms composed of C, H, N, S, and O atoms (these atoms are the only atoms which may appear in the backbone of the divalent radicals), the last being in the form of other linkages. No two hetero atoms may be adjacent within the backbone of the divalent hydrocarbon radical. The value of d is 1, 2, or 3 and n is 1 or 2. R¹⁶ is an oxyhydrocarbyl group, preferably alkoxy, acyl, or acetoxy, having 1 to 8 carbon atoms or a radical of formula (CH₂CH₂O)_kZ in which k is an integer of at least 1 and Z is an aliphatic hydrocarbon radical or less than 10 carbon atoms or hydrogen. An oxyhydrocarbyl group is a hydrocarbon group or hydrocarbon group having no more than one heterostom in the backbone selected from N, S, and O for every two carbon atoms in the backbone, wherein a hydrogen attached to a carbon atom has been replaced with a divalent oxygen atom to form an external bond. For example, starting with CH₃CH₂OCH₃, the oxyhydrocarbyl derivatives include -- O--CH₂CH₂OCH₃. Exemplary hydrocarbyl groups include aromatic groups (e.g., phenyl, naphthyl and benzothienyl) and aliphatic groups (linear, branched, or cyclic) including cyclohexyl, tetrahydrofuryl, dioxanyl, peperidyl, pyrolidinyl, ethoxyethoxy ethyl, etc.). Preferably, this oxyhydrocarbyl group has 1 to 10 carbon atoms.

Examples of useful silanes are vinyltriethoxy silanes, vinyltris(2-methoxyethoxy) silane, vinyltriacetoxy silane, gamma-methacryloxypropyltrimethoxy silane, beta-(3,4,epoxycyclohexyl) ethyltrimethoxy silane and gamma-glycidoxypropyltrimethoxy silane. Other useful silanes of this type are known to the art. Combinations of silanes or additions of silanes and compounds copolymerizable with either functional group of the reactant (silane, epoxy, vinyl or methacryloxy) is contemplated in the practice of this invention. Preferably, however, the composition is free from amino silanes.

Two or more ethylenically unsaturated materials may, of course, be employed in the composition.

The composition preferably comprises at least about 5 parts by weight of the ethylenically unsaturated material. Most preferably it comprises from about 10 to 100 parts by weight of the ethylenically unsaturated material.

A reaction initiator may be incorporated into the coating composition, it comprises at least about 0.1 parts, and preferably from about 1 to 4 parts by weight of the ethylenically unsaturated material.

The nature of the reaction initiator utilized in the composition is dependent upon the ethylenically unsaturated material used. For example, for those ethylenically unsaturated materials which undergo free redical polymerization, suitable initiators are compounds which illberate or generate a free-radical on addition of energy. Such initiators include peroxy, azo, and redox systems each of which are well known and are described in polymerization art, e.g. Chapter 11 of *Photochemistry*, by Calvert and Pltts, John Wiley & Sons (1966). Included among free-radical initiators are the conventional heat activated catalysts such as organic peroxides and organic hydroperoxides. Examples of these catalysts are benzoyl peroxide, tertiary-butyl perbenzoate, cumene hydroperoxide, exobis(isobutyronitrile) and the like.

The preferred catalysts are photopolymerization initiators. Included among such initiators are acyloin and derivatives thereof, such as benzoin, benzoin methyl ether, benzoin ethyl ether, benzoin isopropyl ether, benzoin isobutyl ether, and a-methylbenzoin; diketones such as benzil and diacetyl, etc.; organic sulfides such as diphenyl monosulfide, diphenyl disulfide, decyl phenyl sulfide, and tetramethylthiuram monosulfide; S-acyl dithiocarbamates, such as S-benzoyl-N,N-dimethyldithiocarbamate; phenones such as acetophenone, a,a,a-tribromacetophenone, a,a-diethoxyacetophenone, a,a-dimethoxy-a-phenylacetophenone, o-nitro-a,aa-tribromacetophenone, benzophenone, and p,p'-bis(dimethyliamino)benzophenone; aromatic iodonium and aromatic sulfonium salts, sulfonyl halides such as p-toluenesulfonyl chloride, 1-naphthalenesulfonyl chloride, 1,3-benzenedisulfonyl chloride, 2,4-dinitrobenzenesulfonyl bromide, and p-acetamidobenzenesulfonyl chloride.

The reaction initiator employed with silane and siloxane materials is preferably a metal ester of the formule M— $(OR^{10})_{tt}$, where M is selected from the group consisting of titanium, aluminum, and zirconium, R^{10} is selected from the group consisting of lower alkyl radicals containing from 1 to 8 carbon atoms which are bonded to the oxygen atom, and x is equal to the number of valence bonds of M.

Other reaction initiators, such as acids or bases, may be used with silene and siloxane materials. Representative metal esters which have been found useful in the practice of the present invention include tetraisopropyl titanate ("Tyzor", commercially available from DuPont); tetrabutyl titanate, tetraethyl titanate, tetra 2-ethylhexyl titanate, aluminum isopropoxide, aluminum n-butoxide, and tetraisopropyl zirconate. Each of the metal esters may be utilized either by itself or in combination in the

practice of the present invention. Other useful metal esters of the type described are known to the art. The preferred metal ester is tetraisopropyl titanate.

The coating compositions can include a variety of other ingredients such as compatible monomers and polymers, stabilizers, antioxidants, flexibilizers, colorants (e.g., dyes and pigments), reinforcing fillers, surfactants, flow aids, hardness enhancers (e.g., colloidal silica), and the like. Additionally, solvents may be utilized to facilitate preparation and application of the compositions. Examples of suitable solvents include methyl ethyl ketone, chloroform, methylene chloride and methyl alcohol.

The following examples further illustrate the present invention. Unless otherwise noted, all references to parts are to parts by weight.

Example 1
A coating composition containing the following ingredients was prepared:

Part A		Parts
ČH³CHCH³O(CH³)³	Si(OCH ₂) ₃ (1)	75
8		
O n		
CH2=CCO(CH5)32!(OCH ₃) ₃ ⁽²⁾	25
CH ₃		
Water		50
Part B		•
Methyl Alcohol		65
HCI (Concentrated)		4.7
Tetra isopropyl tita	nate ⁽²⁾	28.4

^m A-187 silane available from Union Carbide Corporation

Part A was prepared by combining the ingredients in a glass beaker at room temperature and stirring until a one-phase product was obtained. Part B was prepared by combining the ingredients in the order listed in a glass beaker at room temperature and stirring. Part A (12 g) and Part B (9 g) were then combined in a glass beaker with stirring at room temperature for about 15 minutes. A minor amount (0.5% by weight) of a fluorochemical surfactant ("FC-430" available from 3M Company) was added to the resultant product and stirring was continued for about 5 minutes.

The concave face of a single vision ophthalmic lens mold was coated with the resulting product by holding the mold at an angle of about 45 degrees from horizontal and painting a small quantity of the composition across the top edge of the concave face along an arc having a central angle of from 140° to 160°. The coating composition flowed downward across the face of the mold and left a thin, uniform, coating on the concave surface. The coating was partially reacted by placing the coated mold about 5 cm from a sunlamp and exposing it to ultraviolet light for two minutes at about 80° to 100°C.

The coated mold was then used to cast a single vision ophthalmic lens blank. Allyl diglycol carbonate monomer ("CR-39" from PPG industries) containing 3% by weight benzoyl peroxide was charged to the mold until the mold was full. The filled mold was placed in a water bath and heated at 65°C for one hour, 73°C for one hour, and then at 85°C for 15 hours. The temperature of the bath was then reduced to 60°C and the resulting hardened lens removed from the mold.

The coating separated completely from the mold face upon removal of the lens from the mold and was firmly bonded to the convex surface of the lens. It could not be removed from the lens when tested according to ASTM D-3359 Tape Adhesion Test. Additionally, the coating exactly replicated the convex surface of the lens, was free from eberrations and had a SEB abrasion resistance of (30—40 psi) 206.7—275.6 kPa. The coating was about 6 microns thick. The coating was subjected to postcuring by heating it for 8 hours at 85°C. After postcuring it had an SEB abrasion resistance of (70 psi) 482.3 kPa.

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⁽²⁾ A-174 silane available from Union Carbide Corporation

^{(3) &}quot;Tyzor" available from DuPont.

Example 2

A poly(ethylenically unsaturated alkyoxyalkyl)heterocyclic compound of the formula

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was prepared as described in U.S. Patent 4,249,011 (Example 1). Sixty parts of the heterocyclic compound and 40 parts of methyl ethyl ketone were combined and mixed until uniform. A polymerization initiator (2% by weight of q,q-dimethoxy-q-phenylacetophenone ("Irgacure" 651 available from Ciba Geigy) was added to the mixture and stirred until a uniform composition was obtained. This composition was applied to the concave surface of a single vision ophthalmic lens mold as described in Example 1 to provide a thin coating thereon.

The methyl ethyl ketone was evaporated from the coating by allowing the coated mold to stand at room temperature for several minutes. The coating was then partially polymerized by exposing it to a 700 watt Hanovia medium pressure mercury arc lamp for 10 seconds at a distance of about 5 cm. The coated die was then used to prepare an ophthalmic lens blank as described in Example 1 using allyl diglycol carbonate monomer and 3% by weight benzoyl peroxide. The lens was cured as described in Example 1.

When the lens was removed from the mold, the coating separated essentially completely from the mold face and remained adhered to the lens surface. The coating was firmly bended to the convex surface of the lens and could not be removed when tested according to ASTM D-3359 Tape Adhesion Test. The coating replicated the convex surface of the lens and was free from aberrations. The coated lens had a SEB abrasion resistance of 275.6—413.4 kPa (40—60 psi) and was 10 to 12 µm thick. The coating was post cured by exposing it to ultraviolet light as described above. After post curing it had an SEB abrasion resistance of (90 psi) 620.1 kPa.

Example 3

A composition was prepared which contained 80 parts of the poly(ethylenically unsaturated alkoxyalkyl) heterocyclic compound described in Example 2, 10 parts of Silane "A-174", 2 parts o.q.dimethoxy-q-phenylacetophenone, and 508 parts by weight of methyl ethyl ketone. The composition was applied to the concave surface of a series of multivision ophthalmic lens molds as described in Example 1. After the solvent was evaporated, the coatings were partially polymerized by exposing them for five seconds to a 700 watt Hanovia medium pressure mercury arc lamp.

The coated dies were then used to prepare multivision ophthalmic lenses using the techniques described in Example 1. After curing and removal from the molds, the coatings separated completely from the mold faces and were firmly bonded to the convex surfaces of the lenses. They could not be removed from the lens surfaces when tested according to ASTM D-3359 Tape Adhesion Test. The exposed surfaces of the coatings exectly replicated the convex surfaces of the lenses and were free from aberrations. The lenses had a SEB abrasion resistance of (30—50 psi) 206.7—344.9 kPa and were free from flow lines and coating buildup at the segment lines of the multivision segments. The coatings were from 4 to 5 μm (microns) thick.

The SEB abresion resistance of the lenses was improved to 75 psl by post exposing them for 2 minutes to ultraviolet light from a 100 watt/in Hanovia Lamp positioned 10 cm from the coating. This did not affect the adhesion of the coatings to the lenses.

Example 4

An ethylenically unsaturated material was prepared which had the formula:

wherein Q was CH₂=CH(CH₃)COO(CH₂)₃, m had a value of between 3 and 8, and p had a value between from 2 to 10. The procedure used was as follows.

A 2 I flask was placed in a heating mantle and fitted with reflux condenser, mechanical stirrer, and thermometer. The flask was charged with 1500 g of amber-colored gamma-methacryloxy-propyltrimethoxysilane (available from Union Carbide as A-174 or from Dow Corning as Z-6030) and 150 g anhydrous sodium carbonate. The flask and contents were heated to about 130°C and maintained at that temperature with stirring for 4 hours. The mixture was cooled to ambient temperature and filtered to remove the insoluble residue. The filtrate was returned to the flask together with about 2.25 g (0.15%) potassium hydroxide pellets. Stirring was maintained for 1 hour without heating as the liquid became very dark brown to black in color. The liquid was decanted and distilled under vacuum at 85 to 110°C and 1 to 5 mm Hg pressure to yield water-white gamma-methacryloxypropyltrimethoxysilane.

A second flask was charged with 150 g of the above purified gamma-methacryloxypropyltrimethoxysilane, 150 ml of tetrahydrofuran (analytical reagent grade) 0.03 g butylated hydroxytoluene (as a stabilizer) and 33.6 g 1 normal hydrochloric acid. The mixture was refluxed for one hour and then evaporated in vacuum (45°C at 5 mm Hg) to provide a residue of water-white clear oligomeric prepolymer of the above formula having a viscosity of about 11,000 cps after sir-purging.

The prepolymer was combined with 2 parts by weight of q,q-dimethoxy-q-phenylacetophenone, warmed slightly, and applied as a thin coating to the concave surface of a multivision ophthalmic-lens mold using the procedures described in Example 1. The coating was partially polymerized by exposing it for 2 seconds to ultraviolet light from a 100 watt/in Hanovia lamp located 10 cm from the coating. The mold was then used to prepare ophthalmic lenses from allyl diglycol carbonate monomer as described in Example 1 except that the lenses were heated in the water bath by slowly raising the temperature of the bath from 60°C to 85°C during a 12 hour period. When the lenses were removed from the mold, the coating separated completely from the mold faces and was firmly bonded to the convex surface of the lenses, it could not be replicated the convex surface of the lenses when tested according to ASTM D-3359 Tape Adhesion test. The coating replicated the convex surface of the lenses and was free from aberrations because the coatings were free from flow lines and coating buildup at the segment lines of the multivision segments. The lenses had an SEB abrasion resistance of (30—45 psi) 206.7—310 kPa.

The abrasion resistance of the lenses was increased to (60 psi) 413.4 kPa by post exposing the coatings for two minutes to ultraviolet from a 100 watt/in Hanovia lamp located 10 cm from the coating. This did not affect the adhesion of the coatings to the lenses.

Example 5
A composition having the following ingredients was prepared:

35		Parts
	Poly(ethylenically unsaturated alkoxyalkyl) heterocyclic compound of Example 2	60
40	Pentaerythritoi Tetraacrylate	40
	"Irgacure" 651	3
45	"FC-431"	0.25
	Methylene Chloride	896.75

This composition was applied to and partially reacted on a series of multivision ophthalmic lens molds so using the methods described in Example 2. Various power levels were utilized during the exposure. The coated molds were then used to prepare ophthalmic lenses from allyl diglycol carbonate as described in Example 1. The coated lenses were tested for residual unsaturation using infrared absorption techniques; SEB abrasion resistance and adhesion. The results of the abrasion resistance and the adhesion were reported as poor, fair, good and excellent. Poor (i.e., unacceptable) abrasion resistance meant that the 55 coating visibly scratched with the application of less than (20 psl) 135 kPa of force. Fair (acceptable) abrasion resistance meant that 20 to 40 psi (135 kPa to 275 kPa of force were required before the coating visibly scratched. Good abrasion resistance means that (40 to 60 psi) 275 kPa to 405 kPa of force were required before the coating visibly scratched. Excellent abrasion resistance means that the coating either did not visibly scratch or visibly scratched only after the application of more than (60 psl) 405 kPa of force. 50 Poor (unacceptable) adhesion means that more than 10% of the coating was removed when tested according to the Tape Adhesion test ASTM D-3359. Good (acceptable) adhesion means that only from 5 to 10% of the coating was removed when tested according to ASTM D-3359. Excellent adhesion means that there was no coating removed when tested according to ASTM D-3359. The results of these tests are set out below.

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	Sample	% Residual Unsaturation	Abresion Resistance	Adhesion
5	A	75	Fair	Excellent
	В	67	Excellent	Excellent
	С	62	Excellent	Excellent
	D	57.1	Excellent	Excellent
	ε	55	Excellent	Excellent
15	F	48.3	Excellent	Fair
	G	43	Excellent	Fair
	н	36.1	Excellent	Poor
20	ı	30.6	Excellent	Poor

Example 6

Coating composition prepared as described in Examples 2, 3 and 5 were applied to the convex surfaces of ophthalmic lens molds using the coating technique described in Example 1. The coated molds were allowed to air dry at ambient room temperature for about one minute and then separate molds were partially cured by exposing them to a 300 watt/in Hanovia medium pressure mercury lamp. The coatings were then tested for SEB abrasion resistance. The results are set out below.

30	Coating Sol'n from Ex	% Residual Unsaturation	Abrasion Resistance (psl)
	. 2	30	<7.5 <51.68 kPa
25	2	60	<7.5 <51.68 kPa
	2	80	<7.5 <51.68 kPa
40	3	30	<7.5 <51.68 kPa
•	3	60	<7.5 <51.68 kPa
	3	90	<7.5 <51.68 kPa
45	5	30	<7.5 <61.68 kPa
	5	60	<7.5 <51.68 kPa
50	5	90 (<7.5 <51.68 kPa

The coatings each had scratches so numerous that they became opaque.

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Example 7
A coating composition having the following ingredients was prepared:

	<u>.</u>	Parts
5	CH2CHCH2O(CH2)3SI(OCH3)3	14.08
	0	
10	CH ₂ =CCO(CH ₂) ₃ Si(OCH ₃) ₃	4.7
	ĊH₃	
	Delonized water	7.74
15	Methyl alcohol	6.71
	HCI (37%)	.96
20 .	Tetraisopropyl Titanate	5.64
	Methyl ethyl ketone	59.92
	Fluorochemical surfactant (FC-430)	0.24

Clean glass and metal dies were flow coated with the above composition using the technique described in Example 1. The coated dies were allowed to stand at room temperature for about two minutes and then partially reacted for two minutes at 100°C using an infrared heat lamp. Ophthalmic lenses were then cast from sikyl diglycol carbonate using the molds. The lenses were hardened by heating them at 48—85°C for 4½ hours. The lenses were removed from the mold, given a post cure at 90°C for four hours and tested for SEB abrasion resistance and adhesion. The coating had an SEB abrasion resistance of 482.3 kPa (70 psi) and 100% adhesion to the lens.

Example 8
Two separate costing solutions having the following ingredients were prepared:

35	Solution A	Parts
	CH2CHCH2O(CH2)3SI(OCH3)3	18.78
40	o -	
	Deionized water	7.74
	Methyl Alcohol	6.71
45	HCI (37%)	0.96
	Tetraisopropyl titanate	5.64
Fa	Methyl ethyl ketone	59.92
50	Fluorocarbon surfactant (FC-430)	0.24
	Solution B	Parts
55	Poly(ethylenically unsaturated alkoxy- alkyl)heterocyclic compound of Examp	ole 2 6
	Pentaerythritol Tetrascrylate	4
. 60	Photoinitiator (Irgacure® 651)	0.3
	Fluorocarbon surfactant (FC-431)	0.03
65	Methylene Chloride	89.67

Solution A was flow costed directly onto primed trifocal lenses and then partially cured using a 200 watt/in medium pressure lamp (1 pass, 0.25 joule/cm²). The lenses were then post cured at 80°C for 12 hours.

Trifocal lenses coated with solution B were prepared using the techniques described in Example 1 except that the coated molds were partially cured using a 200 watt/in medium pressure lamp and exposing the molds to only one pass at 0.25 joules/cm². The filled molds were heated to 48—85°C for seven hours to cure the lenses. The cured lenses were removed from the molds and post cured using a 300 watt/in medium pressure lamp and exposing them to four passes at 1 joule/cm².

A Ronchi Graph was used to test the aberration of the coated surfaces. The trifocal lenses coated with solution A all exhibited aberrations around the segments while the trifocal lenses coated with Solution B did not exhibit aberrations.

Example 9

Four coated ophthalmic lenses were prepared using the coating solution of Example 5. The coating solutions were applied and partially cured as described in Example 1 except that a level of 0.25 joules/cm² were used for partial curing. After the lenses had been hardened, the coatings were exposed to 4 joules/cm². The coatings had 100% adhesion to the lenses.

The coated lenses were dipped into a 93°C water bath for 10 minutes, removed and immediately dipped into a 2°C ice/water bath for 10 minutes. This cycle was repeated 5 times.

The coatings were then tested for coating adhesion and found to have 100% adhesion to the lenses.

Claims

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1. A method of providing a thin, optically clear coating to at least one optical surface of an ophthalmic 25 lens comprising the steps of:

(i) applying a layer of a composition comprising a material containing reactive ethylenically unsaturated groups to at least a first face of a mold used to manufacture said ophthalmic lens, wherein said face imparts a desired optical configuration to a first optical surface of said ophthalmic lens;

(ii) reacting said composition to a degree that said composition forms a film that is dry, non-abrasion resistant, at least weakly adhered to said first face, and exactly replicates said first face in an aberration-free manner such that said composition exhibits a degree of unsaturation in the range of 40% to 90% of the unsaturation it possessed prior to reaction;

(iii) filling said mold with an organic liquid material capable of hardening to a solid, room temperature-

stable state; and

(iv) hardening said organic liquid material so as to form said ophthalmic lens, intimately bond said film to the hardened organic material, and adhere said film to said optical surface of said hardened organic material more firmly than it adheres to said face, and

(v) post reacting said composition after said organic liquid has hardened to harden said composition.

2. A method as claimed in claim 1 wherein said material is a poly(ethylenically unsaturated alkoxyalkyl) compound of the formula A1—Z—A2 wherein

(a) A¹ and A² are independently groups having terminal ethylenic unsaturation said groups having the formula:

in which

(1) R¹ is selected from hydrogen or methyl
(2) R—O— is a monovalent residue of an aliphatic terminally unsaturated primary alcohol, ROH, where

R is selected from the formulae:

ss and

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wherein E is selected from the formulae:

65 a and c are independently integers of from 1 to 6, b is zero or an integer of from 1 to 6, R4 is selected from

hydrogen or methyl, R⁵ is an aliphatic group having 1 to 15 carbon atoms which may be interrupted with up to two groups selected from the class consisting of ether or oxygen or

Rs having a valence of m+1 wherein m is an integer of 1 to 5,

(3) R2 is selected from the group of hydrogen,

wherein R⁶ is selected from alkyl and alkenyl groups having up to 5 carbon atoms, R⁷ is an aliphatic group of up to 8 carbon atoms or an aromatic group of up to 8 carbon atoms,

(4) R3 is an alkylene group having 1 to 6 carbon atoms and up to one catenary oxygen in the group, and

(b) Z is a heterocyclic group of the formula:

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wherein X is a divalent group required to complete a 5-membered heterocyclic ring and is selected from the

wherein R^{a} , R^{a} , R^{1a} , and R^{1} are independently selected from hydrogen, alkyl groups of 1 to 4 carbon atoms, cycloalkyl groups of 3 to 6 carbon atoms, and phenyl groups, and A3 is the same as A1, and wherein said non-abrasion-resistant film is reacted in step (v) to harden it to an SEB abrasion resistance of at least 137.9

3. A method according to claim 1 or claim 2 including the further step of removing said lens from said mold characterised in that said film adheres to said first optical surface and separates essentially completely from said mold face.

4. A method according to any preceding claim characterised in that seld composition is partially reacted by exposing it to a temperature in the range of from 35°C to 100°C for from 1 to 10 minutes.

5. A method according to any preceding claim characterised in that said face of said mold is positioned at an angle of from the horizontal in the range of about 45° to 60° prior to applying said composition thereto.

6. A method according to claim 5 characterised in that said composition is applied to the top edge of said first face so that said composition covers an arc having a central angle in the range of from about 140° to 160°, and allowed to flow across said face of said mold.

7. A method according to any preceding claim characterised in that said composition is reacted by exposure to ultraviolet light.

8. A method according to any of claims 1 to 6 characterised in that said composition is reacted by

exposure to electron beam radiation.

9. A method according to any preceding claim characterised in that said coating is reacted until it exhibits a degree of unsaturation in the range of from 40% to 75% of the unsaturation it possessed prior to

10. A method according to claim 9 characterised in that said coating is reacted until it exhibits a degree

of unsaturation in the range of from 55% to 70%.

11. A method according to any preceding claim comprising the further steps of applying a thin layer of said composition to a second face of said mold, characterised in that said second face forms a desired optical configuration on a second surface of said ophthalmic device, and partially reacting said composition on said second face.

12. A method according to any preceding claim characterised in that said ophthalmic lens comprises a multifocal spectacle lens.

13. A method according to any preceding claim characterised in that said composition further includes

14. A lens coated according to any preceding claim characterised in that said composition further includes one or more materials selected from the group consisting of polymers, stabilizers, anti-oxidants, flexibilizers, colorants, fillers, surfactants, flow aids, and hardness enhancers.

Patentansprüche

1. Verfahren zum Ausbilden eines dünnen, optisch klaren Überzuges auf mindestens einer optischen Fläche einer Augenlinse, mit folgenden Schritten:

(i) auf mindestens eine erste Fläche einer zum Herstellen der Augenlinse verwendeten Form word eine Schicht aus einer Zusammensetzung aufgetragen, die mindestens zum Teil aus einem Material besteht, das ethylenisch ungesättigte, reaktionsfählge Gruppen enthält, wobei die genannte Fläche einer ersten optischen Fläche der Augenlinse eine gewünschte optische Form erteilt;

(II) die genannte Zusammensetzung wird derart umgesetzt, daß die genannte Zusammensetzung eine Schicht bildet, die trocken und nicht abreibfest ist, an der genannten Fläche mindestens schwach haftet und einen keinen abbildungsfehlerfreien Abdruck der ersten Fläche darstellt, wobei die Zusammensetzung eine Ungesättigtheit im Bereich von 40 bis 90% der Ungesättigtheit besitzt, die sie vor der Umsetzung besaß;

(iii) die Form wird mit einem organischen flüssigen Material gefüllt, das geeignet ist, in einen festen,

bei Zimmertemperatur beständigen Zustand zu erhärten; und

(iv) durch Härten des organischen flüssigen Materials wird die Augenlinse gebildet, die genannte Schicht mit dem gehärteten organischen Material in satter Anlage daran stoffschlüssig verbunden und die Schicht mit der optischen Fläche des gehärteten organischen Materials fester verldebt als mit der genannten Fläche, und

(v) nach dem Härten der organischen Flüssigkeit wird durch eine Nachreaktion der Zusammensetzung

20 diese gehärtet.

2. Verfahren nach Anspruch 1, dadurch gekennzeichnet, daß das genannte Material ein Poly(ethylenisch ungesättigte Alkoxyalkyl)-Verbindung der Formel A¹—Z—A² ist, in der

(a) A' und A' unabhängig voneinander endständig ethylenisch ungesättigte Gruppen der Formel

30 sind, in der

(1) R1 aus Wesserstoff und Methyl ausgewählt ist:

(2) R—O— ein einwertiger Rest eines endständig ungesättigten eliphatischen ungesättigten Alkohols ROH ist, wobei R aus der Formeln

und

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ausgewählt ist, E aus den Formein

ausgewählt ist.

a und c unabhängig voneinander ganze Zahlen von 1 bis 6 sind, b null oder eine ganze Zahl von 1 bis 6 ist, R⁴ aus Wasserstoff und Methyl ausgewählt ist, R⁵ eine eliphatische Gruppe ist, die 1 bis 15 Kohlenstoffatome besitzt und mit bis zu zwei Gruppen unterbrochen sein kann, die aus der Gruppe ausgewählt sind, die aus Äther, Sauerstoff und den Gruppen

ausgewähit sind,

R⁶ eine Wertigkeit von m+1 hat, wobei m eine ganze Zahl von 1 bis 5 ist.

(3) R² aus der Gruppe Wasserstoff

ausgewählt ist, wobel R^e aus den Alkyl- und Alkenylgruppen mit bis zu 5 Kohlenstoffatomen ausgewählt ist, R7 eine aliphatische Gruppe mit bis zu 8 Kohlenstoffatomen oder eine aromatische Gruppe mit bis zu 8 Kohlenstoffstomen ist.

(4) R³ eine Alkenylengruppe mit bis 6 Kohlenstoffatomen und bis zu einem Kettensauerstoffatom in der Gruppe ist und

(b) Z eine heterozyklische Gruppe der Formel

ist, in der X eine zweiwertige Gruppe ist, die zum Vervollständigen eines 5-gliedrigen heterozyklischen Ringes erforderlich ist und aus der Gruppe

ausgewählt ist, in der R⁸, R⁹, R¹⁰ und R¹¹ unabhängig voneinander aus Wasserstoff, den Alkylgruppen mit 1 bis 4 Kohlenstoffatomen, den Cycloalkylgruppen mit 3 bis 6 Kohlenstoffatomen und den Phenylgruppen ausgewählt sind, A³ dasselbe ist wie A¹, und

die nicht abreibfeste Schicht durch die Reaktion im Schritt (v) zu einer SEB-Abriebfestigkeit von

mindestens 137,9 kPa gehärtet wird. 3. Verfahren nach Anspruch 1 oder 2 mit einem weiteren Schritt, in dem die Augenlinse aus der Form hersusgenommen wird, dadurch gekennzeichnet, daß die genannte Schicht an der ersten optischen Fläche haften und sich von der genannten Fläche der Form im wesentlichen vollständig ablöst.

4. Verfahren nach einem der vorhergehenden Ansprüche, dadurch gekennzeichnet, daß die Zusammensetzung zu ihrer teilweisen Umsetzung 1 bis 10 min lang auf einer Temperatur im Bereich von

35 bis 100°C gehalten wird. 5. Verfahren nach einem der vorhergehenden Ansprüche, dadurch gekennzeichnet, daß die genannte Fläche der Form vor dem Auftragen der Zusammensetzung auf diese Fläche unter einem Winkel von etwa

45 bis 60 Grad zur Horizonteien angeordnet wird. 6. Verfahren nach Anspruch 5, dadurch gekennzeichnet, daß die Zusammensetzung auf den oberen Rand der ersten Häche derart aufgetragen wird, daß die Zusammensetzung einen Bogen mit einem Zentriwinkel im Bereich von etwa 140 bis 160 Grad bedeckt, und daß ihr gestattet wird, über die genannte Häche der Form zu fließen.

7. Verfahren nach einem der vorhergehenden Ansprüche, dadurch gekennzeichnet, daß die

Zusammensetzung zu ihrer Reaktion Ultraviolettilicht ausgesetzt wird.

8. Verfahren nach einem der Ansprüche 1 bis 6, dadurch gekennzeichnet, daß die Zusammensetzung

zu Ihrer Umsetzung einer Elektronenstrahlbestrahlung ausgesetzt wird.

9. Verfahren nach einem der vorhergehenden Ansprüche, dadurch gekennzeichnet, daß der Überzug umgesetzt wird, bis seine Ungesättigtheit im Bereich von 40 bis 75% seiner Ungesättigtheit vor der Umsetzung beträgt.

10. Verfahren nach Anspruch 9, dadurch gekennzelchnet, daß der Überzug umgesetzt wird, bis selne Ungesättigtheit 55 bis 70% beträgt.

11. Verfahren nach einem der vorhergehenden Ansprüche mit weiteren Schritten, in denen eine dunne Schicht aus der Zusammensetzung auf eine zweite Fläche der Form aufgetragen wird, dadurch gekennzeichnet, daß diese zweite Fläche einer zweiten Fläche der Augenlinse eine gewünschte optische Form erteilt, und daß die Zusammensetzung auf der zweiten Fläche tellumgesetzt wird.

12. Verfahren nach einem der vorhergehenden Ansprüche, dadurch gekennzeichnet, daß die Augenlinse ein multifokales Brillenglas ist.

13. Verfahren nach einem der vorhergehenden Ansprüche, dadurch gekennzeichnet, daß die Zusammensetzung ferner einen Reaktionsinitiator enthält.

14. Linse, die nach einem der vorhergehenden Ansprüche mit einem Überzug versehen worden ist, dadurch gekennzeichnet, daß die Zusammensetzung ferner ein oder mehrere Materialien enthält, die aus der Gruppe ausgewählt sind, die aus den Polymeren, Stabilisatoren, Antioxidantien, Flexibilisierungsmitteln, färbenden Stoffen, Füllstoffen, Tensiden, Fließhilfen und Härteverbesserern

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Revendications

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1. Procédé de production d'un revêtement mince optiquement clair sur au moins une surface optique

d'une lentille ophtalmique, qui comprend les étapes consistant:

(i) à appliquer une couche d'une composition comprenant une matière contenant des groupes réactifs à non-saturation éthylénique sur au moins une première face d'un moule utilisé pour fabriquer ladite lentille ophtaimique, ladite face conférant une configuration optique désirée à une première surface optique de ladite lentille ophtelmique;

(ii) à faire réagir ladite composition jusqu'à un degré tel qu'elle forme un film qui soit sec, non résistant 10 à l'abrasion, qui adhère au moins légèrement à ladité première face et qui reproduise exactement cette première face d'une manière dépourvue d'aberrations afin que ladite composition présente un degré de non-saturation dans l'intervalle de 40 à 90% de la non-saturation qu'il possédait avant la réaction;

(iii) à remplir le moule avec une matière líquide organique capable de durcir jusqu'à un état solide

stable à la température ambiante; et

(iv) à faire durdr la matière organique liquide de manière à former ladite lentille ophtalmique, à créer une liaison intime entre le film et la matière organique durcie et à faire adhérer le film à la surface optique de la matière organique durcie plus fermement qu'il n'adhère à ladite face, et

(v) à effectuer la post-réaction de ladite composition après le durcissement du liquide organique pour

faire durcir cette composition.

2. Procédé selon la revendication 1, dans lequel la matière est un composé poly(alkoxyalkylique à nonsaturation éthylénique) de formule A1-Z-A2 dans laquelle

(a) A¹ et A² représentent indépendamment des groupes à non-saturation éthylénique terminale, lesdits

groupes répondant à la formule:

dans laquelle

(1) R1 est choisi entre l'hydrogène et le groupe méthyle, (2) R—O— est un résidu monovalent d'un alcool primaire aliphatique ROH à non-saturation terminale, dans lequel R est choisi entre les formules:

où E est choisi entre les formules:

a et c représentent indépendamment des nombres entiers de 1 à 6, b est égal à zéro ou à un nombre entier de 1 à 6, R⁴ est choisi entre l'hydrogène et le groupe méthyle, R⁶ est un groupe aliphatique ayant 1 à 15 atomes de carbone qui peut être interrompu avec jusqu'à deux groupes choisis dans la classe comprenant de l'oxygène d'éther ou des groupes

R^a ayant une valence m+1 dans laquelle m est un nombre entier de 1 à 5,

(3) R2 est choisi entre l'hydrogène et les groupes

où R^e est choisi entre des groupes alkyle et alcényle ayant jusqu'à 5 atomes de carbone, R⁷ est un groupe aliphatique ayant jusqu'à 8 atomes de carbone ou un groupe aromatique ayant jusqu'à 8 atomes de

(4) R³ est un groupe alkylène ayant 1 à 6 atomes de carbone avec jusqu'à un atome d'oxygène dans la chaîne du groupe, et (b) Z est un groupe hétérocyclique de formule:

dans laquelle X est un groupe divalent nécessaire pour compléter un noyau hétérocyclique pentagonal et est choisi entre les groupes de formules:

dans lesqualles R*, R*, R** et R** sont choisis indépendamment entre l'hydrogène, des groupes alkyle ayant 1 à 4 atomes de carbone, des groupes cycloalkyle ayant 3 à 6 atomes de carbone et des groupes phényle, et A³ est identique à A¹, et ledit film non résistant à l'abrasion est amené à réagir dans l'étape (v) de manière qu'il durcissa jusqu'à ce qu'il acquière une résistance à l'abrasion SEB d'au moins 137,9 kPa (20 lb/in²).

3. Procédé sulvant la revendication 1 ou la revendication 2, comprenant en outre l'opération qui consiste à démouler ladite lentille, caractérisé en ce que le film adhère à la première surface optique et se détache essentiellament en totalité de ladite face du moule.

4. Procédé suivant l'une quelconque des revendications précédentes, caractérisé en ce que la composition est amenée à réagir partiellement par exposition à une température comprise dans la plage de

35 à 100°C pandant 1 à 10 minutes. 5. Procédé suivant l'une quelconque des revendications précédentes, caractérisé en ce que la face du moule est disposée selon un angle par rapport à l'horizontale compris dans la plage d'environ 45 à 60°

avant l'application de ladite composition. 6. Procédé suivant la revendication 5, caractérisé en ca que la composition est appliquée au bord supérieur de la première face de manière que cette composition couvre un arc ayant un angle au centre compris entre environ 140 et 160°, et on la laisse s'écouler en travers de ladite face du moule.

7. Procédé suivant l'une quelconque des revendications précédentes, caractérisé en ce que la composition est amenée à réagir par exposition à la lumière ultraviolette.

8. Procédé suivant l'une que conque des revendications 1 à 6, caractérisé en ce que la composition est amenée à réagir par exposition à un falsceau d'électrons.

9. Procédé suivant l'une quelconque des revendications précédentes, caractérisé en ce que le revêtement est amené à réagir jusqu'a ce qu'il présente un degré de non-saturation dans l'intervalle de 40 à 75% de la non-saturation qu'il possédait avant la réaction.

10. Procédé sulvant la revendication 9, caractérisé en ce que le revêtement est amené à réagir jusqu'à

ce qu'il présente un degré de non-saturation dans la plage de 55 à 70%.

11. Procédé suivant l'une quelconque des revendications précédentes, qui comprend les opérations complémentaires consistant à appliquer une mince couche de composition à une seconde face du moule, caractérisé en ca que la seconde face du moule forme une configuration optique désirée sur une seconde surface du dispositif ophtalmique, et ladite composition est amenée à réagir partiellement sur ladite seconde face.

12. Procédé suivant l'une quelconque des revendications précédentes, caractérisé en ce que la lentille ophtalmique est un verre de lunette multi-focal.

13. Procédé suivant l'une quelconque des revendications précédentes, caractérisé en ce que la

composition comprend en outre un initiateur de réaction. 14. Lentille revêtue suivant l'une quelconque des revendications précédentes, caractérisée en ce que la composition contient en outre une ou plusieurs matières choisies entre de polymères, des agents stabilisants, des anti-oxydants, des assouplissants, des colorants, des charges, des surfactants, des adjuvants d'écoulement et des agents améliorant le durcissement.

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